1) RCHO, 1 or 2 equiv DMF, room temp to 70°C, 0.5 to 2 hours 2) NaBH3CN, 1 to 2 equiv 2) NaBH3CN, 1 to 2 equiv roam temp to 70°C, 1 to 2 hours

Two mono-N-alkyl and one di-N-alkyl vancomycins

taining the additional mass due to alkylation on the amino group of vancosamine. mycin substituted on vancosamine gives vancosaminyl-O-glucose and vancosamine fragments conleucine fragments show increased mass corresponding to the alkyl residue. The mono-N-alkyl vancocosamine fragments, whereas the devancosamine vancomycin, aglucovancomycin, and N-methyl

SAR of N-Alkyl Vancomycins

The first derivatives prepared were the N-decyl vancomycins with an aliphatic straight chain

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Fig. 2. FAB-MS fragmentation pattern of N-alkyl vancomycins.

	305 + R = Vancosamine + glucose + R	159 + R = Vancosamine + R + oxygen	143+R = Vancosamine+R	1,143 = Aglucovancomycin + H	1,305 = Devancosamine vancomycin + H	M+H=Vancomycin+R+H=1,447+R+H	R on vancosamine
CH, -1 -1 -1 -1 -1	: 3 - 1 	99 + R = 13C \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	£ 0.	1,143+R=Aglucovancomycin+R+H	1,305+R = Devancosamine vancomycin+R+H	M+H=Vancomycin+R+H=1,447+R+H	R on leucine

extensive SAR of N-alkyl vancomycins, and over eighty derivatives were prepared and evaluated. in vivo, and shows longer elimination half-life in rats. Encouraged by this result, we undertook an activities of the N-decyl vancomycins with the corresponding N-decanoyl vancomycins shows that similar to the naturally occurring N-acylamido glycopeptides. 1-0) A comparison of the antibacterial N-decyl vancomycin 5 is more active in vitro than the parent vancomycin, equivalent to vancomycin the C₁₀ alkyl analogs are more active than the corresponding alkanoyl series. Furthermore, the mono-

the SAR of the N-acyl vancomycin series, 8) the general trend is that the N-alkyl derivatives substituted on vancosamine are more active than those substituted on N-methylleucine, and both mono-substituted five aromatic N-alkyl vancomycins were prepared and their antibacterial activity compared. As in vancomycins are more active than the corresponding di-N-alkyl vancomycins. A series of the two mono-N-alkyl and one di-N-alkyl derivatives belonging to nine aliphatic and

active of the three derivatives, the reaction conditions for compounds described on Table 3 were ad-Justed so that the most active mono-N-alkyl derivative was the major product of the reaction. The other product for the ten series of compounds in Table 3 were either the other mono-N-alkylated derivative or the di-N-alkyl compound. Having established that the mono-N-alkyl vancomycins substituted on vancosamine are the most

stituted on vancosamine were determined. The pharmacokinetics of the most active derivatives and The in vitro and in vivo antibacterial activity of all thirty-eight mono-N-alkyl vancomycins sub-